



THE UNIVERSITY *of* EDINBURGH

Edinburgh Research Explorer

3D spatial exploration by *E. coli* echoes motor temporal variability

Citation for published version:

Figueroa-Morales, N, Soto, R, Junot, G, Darnige, T, Douarche, C, Martinez, V, Lindner, A & Clément, E 2020, '3D spatial exploration by *E. coli* echoes motor temporal variability', *Physical Review X*, vol. 10, no. 2, 021004. <https://doi.org/10.1103/PhysRevX.10.021004>

Digital Object Identifier (DOI):

[10.1103/PhysRevX.10.021004](https://doi.org/10.1103/PhysRevX.10.021004)

Link:

[Link to publication record in Edinburgh Research Explorer](#)

Document Version:

Peer reviewed version

Published In:

Physical Review X

General rights

Copyright for the publications made accessible via the Edinburgh Research Explorer is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy

The University of Edinburgh has made every reasonable effort to ensure that Edinburgh Research Explorer content complies with UK legislation. If you believe that the public display of this file breaches copyright please contact openaccess@ed.ac.uk providing details, and we will remove access to the work immediately and investigate your claim.



Important Notice to Authors

No further publication processing will occur until we receive your response to this proof.

Attached is a PDF proof of your forthcoming article in *Physical Review X*. The article accession code is XU10353.

Your paper has been assigned the following Subject Area labels: Biological Physics, Interdisciplinary Physics, Soft Matter
Your paper will be in the following section of the journal: RESEARCH ARTICLES

Please note that as part of the production process, APS converts all articles, regardless of their original source, into standardized XML that in turn is used to create the PDF and online versions of the article as well as to populate third-party systems such as Portico, Crossref, and Web of Science. We share our authors' high expectations for the fidelity of the conversion into XML and for the accuracy and appearance of the final, formatted PDF. This process works exceptionally well for the vast majority of articles; however, please check carefully all key elements of your PDF proof, particularly any equations or tables.


Figures submitted electronically as separate files containing color appear in color in the journal.

Specific Questions and Comments to Address for This Paper

The numbered items below correspond to numbers in the margin of the proof pages pinpointing the source of the question and/or comment. The numbers will be removed from the margins prior to publication.

- 1 Third proof:** Please note that only funding organization names are allowed in “Funding Information” section, program number has been fixed as per PRX style.
- 2 Third proof:** Please check very closely to confirm all of the corrections have been incorporated properly throughout this Letter and also approve **the Letter is ready to be published in its current form.**

ORCIDs

Please follow any ORCID links () after the authors' names and verify that they point to the appropriate record for each author.

Funding Information

Information about an article's funding sources is now submitted to Crossref to help you comply with current or future funding agency mandates. Crossref's Open Funder Registry (<https://www.crossref.org/services/funder-registry/>) is the definitive registry of funding agencies. Please ensure that your acknowledgments include all sources of funding for your article following any requirements of your funding sources. Where possible, please include grant and award ids. Please carefully check the following funder information we have already extracted from your article and ensure its accuracy and completeness:

- Agence Nationale de la Recherche, FundRef ID <http://dx.doi.org/10.13039/501100001665> (FR/Republic of France)
- Franco-Chilean Ecosud Collaborative Program
- Pierre-Gilles de Gennes Foundation
- H2020 European Research Council, FundRef ID <http://dx.doi.org/10.13039/100010663> (EU/European Union)
- Fondo Nacional de Desarrollo Científico y Tecnológico, FundRef ID <http://dx.doi.org/10.13039/501100002850> (CL/Republic of Chile)
- Ministerio de Economía, Fomento y Turismo, FundRef ID <http://dx.doi.org/10.13039/501100005886> (CL/Republic of Chile)

- H2020 European Research Council, FundRef ID <http://dx.doi.org/10.13039/100010663> (EU/European Union)
- École Supérieure de Physique et de Chimie Industrielles de la Ville de Paris, FundRef ID <http://dx.doi.org/10.13039/501100003068> (FR/Republic of France)
- Laboratoire International Associé “Matière: Structure et dynamique” (LIA-MSD)
- Centre National de la Recherche Scientifique, FundRef ID <http://dx.doi.org/10.13039/501100004794> (FR/Republic of France)
- Institut Pierre-Gilles de Gennes (Équipement d'Excellence, “Investissements d'avenir”)

Other Items to Check

- Please note that the original manuscript has been converted to XML prior to the creation of the PDF proof, as described above. Please carefully check all key elements of the paper, particularly the equations and tabular data.
- Please ignore the text of the Popular Summary, it is not the final version. An APS science writer will work with you separately to edit the summary.
- Title: Please check; be mindful that the title may have been changed during the peer-review process.
- Author list: Please make sure all authors are presented, in the appropriate order, and that all names are spelled correctly.
- Please make sure you have inserted a byline footnote containing the email address for the corresponding author, if desired. Please note that this is not inserted automatically by this journal.
- Affiliations: Please check to be sure the institution names are spelled correctly and attributed to the appropriate author(s).
- Receipt date: Please confirm accuracy.
- Acknowledgments: Please be sure to appropriately acknowledge all funding sources.
- Hyphenation: Please note hyphens may have been inserted in word pairs that function as adjectives when they occur before a noun, as in “x-ray diffraction,” “4-mm-long gas cell,” and “*R*-matrix theory.” However, hyphens are deleted from word pairs when they are not used as adjectives before nouns, as in “emission by x rays,” “was 4 mm in length,” and “the *R* matrix is tested.”
Note also that Physical Review follows U.S. English guidelines in that hyphens are not used after prefixes or before suffixes: superresolution, quasiequilibrium, nanoprecipitates, resonancelike, clockwise.
- Please check that your figures are accurate and sized properly. Make sure all labeling is sufficiently legible. Figure quality in this proof is representative of the quality to be used in the online journal. To achieve manageable file size for online delivery, some compression and downsampling of figures may have occurred. Fine details may have become somewhat fuzzy, especially in color figures. Figures to be published in color online will appear in color on these proofs if viewed on a color monitor or printed on a color printer.
- Overall, please proofread the entire *formatted* article very carefully. The redlined PDF should be used as a guide to see changes that were made during copyediting. However, note that some changes to math and/or layout may not be indicated.

Ways to Respond

- **Web:** If you accessed this proof online, follow the instructions on the web page to submit corrections.
- **Email:** Send corrections to aps-robot@luminad.com. Include the accession code XU10353 in the subject line.
- **Fax:** Return this proof with corrections to +1.855.808.3897.

If You Need to Call Us

You may leave a voicemail message at +1.855.808.3897. Please reference the accession code and the first author of your article in your voicemail message. We will respond to you via email.

Popular Summary

Bacterial motion determines the changing structure of microbial communities and controls infection spreading as well as microbiota organization in ecosystems. *E. coli* bacteria explore their environment using a “run-and-tumble” strategy: a sequence of straight paths (runs) and sudden changes in swimming direction (tumbles) that happen when motors driving the cell’s tail-like flagellum change rotation direction for a short time. While this random walk is classically described as a process with a single characteristic run time, the flagellar motor rotation switching, responsible for reorientations, displays a wide distribution of times. To address this paradox, we built a 3D tracking microscope suited to follow swimming bacteria for as long as tens of minutes.

Our results reconcile individual motor rotation and bacterial spatial exploration in three dimensions. We reveal a continuous variation of exploration “moods” for individual bacteria, characterized by periods of frequent directional changes alternating with periods of persistent swimming. The dynamics can be explained by important fluctuations in the number of certain proteins inside the cell that are responsible for the motor switching.

Future research will address the importance of these realistic run-and-tumble statistics in the macroscopic transport of bacteria. Bacterial persistent swimming may help to explain the onset of medical emergencies as well as bacterial anomalous transport in confined environments, such as narrow capillaries and porous media. This knowledge could be relevant to emerging technologies for targeted drug delivery or for understanding the spreading of biocontaminants in soils.

3D Spatial Exploration by *E. coli* Echoes Motor Temporal Variability

Nuris Figueroa-Morales^{1,2}, Rodrigo Soto³, Gaspard Junot¹, Thierry Darnige¹, Carine Douarche⁴,
Vincent A. Martinez⁵, Anke Lindner¹, and Éric Clément^{1,*}

¹PMMH, UMR 7636 CNRS, ESPCI-Paris, PSL Research University, Sorbonne Université, Université de Paris, 7-9 quai Saint-Bernard, 75005 Paris, France

²Department of Biomedical Engineering, The Pennsylvania State University, University Park, Pennsylvania 16802, USA

³Departamento de Física, FCFM, Universidad de Chile, Santiago 8370448, Chile

⁴Université Paris-Saclay, CNRS, FAST, 91405 Orsay, France

⁵SUPA and School of Physics & Astronomy, The University of Edinburgh, Edinburgh EH9 3FD, United Kingdom

(Received 6 January 2020; accepted 26 February 2020)

Unraveling bacterial strategies for spatial exploration is crucial for understanding the complexity in the organization of life. Bacterial motility determines the spatiotemporal structure of microbial and controls infection spreading and the microbiota organization in guts or in soils. Most theoretical approaches for modeling bacterial transport rely on their run-and-tumble motion. For *Escherichia coli*, the run-time distribution is reported to follow a Poisson process with a single characteristic time related to the rotational switching of the flagellar motors. However, direct measurements on flagellar motors show heavy-tailed distributions of rotation times stemming from the intrinsic noise in the chemotactic mechanism. Currently, there is no direct experimental evidence that the stochasticity in the chemotactic machinery affects the macroscopic motility of bacteria. In stark contrast with the accepted vision of run and tumble, here we report a large behavioral variability of wild-type *E. coli*, revealed in their three-dimensional trajectories. At short observation times, a large distribution of run times is measured on a population and attributed to the slow fluctuations of a signaling protein triggering the flagellar motor reversal. Over long times, individual bacteria undergo significant changes in motility. We demonstrate that such a large distribution of run times introduces measurement biases in most practical situations. Our results reconcile the notorious conundrum between run-time observations and motor-switching statistics. We finally propose that statistical modeling of transport properties, currently undertaken in the emerging framework of active matter studies, should be reconsidered under the scope of this large variability of motility features.

DOI:

Subject Areas: Biological Physics,
Interdisciplinary Physics, Soft Matter

I. INTRODUCTION

The *run-and-tumble* (R&T) strategy developed by bacteria for exploring their environment is a cornerstone of quantitative modeling of bacterial transport. In this paradigm, bacteria swim straight during a *run time*, undergo a reorientation process during a *tumbling time*, and pursue thereafter the next run in a different direction. The now standard vision of the R&T strategy was established in the 1970s for swimming *Escherichia coli* by Berg and Brown [1,2], based on 3D trajectories obtained via a Lagrangian

tracking technique. They proposed that an adapted bacterium would perform, over long times, an isotropic random walk composed of the run-and-tumble phases, both distributed in time as a Poisson process [1–5]. For quantitative analysis, the run-time and tumble-time distributions are often taken as Poisson processes with typical values $\bar{\tau}_{\text{run}} \sim 1$ s and $\bar{\tau}_{\text{tumble}} \sim 1/10$ s [2,6]. These values change in the presence of chemical gradients, leading to a biased random walk known as chemotaxis.

Alongside the relevance of this result in the context of biology, medicine, or ecology, fluids laden with motile bacteria have become an epitome for active matter, where the organization of active particles recently led scientists to revisit many concepts of out-of-equilibrium statistical physics [7–10]. Suspensions of motile bacteria are systems of choice for these studies [11], and many original phenomena such as anti-Fick's law migration [12], collective motion [13], viscosity reduction [14–16], enhanced

*eric.clement@upmc.fr

Published by the American Physical Society under the terms of the [Creative Commons Attribution 4.0 International license](#). Further distribution of this work must maintain attribution to the author(s) and the published article's title, journal citation, and DOI.

diffusion [7], or motion rectification [17–20] have been discovered. Most recent theoretical studies on active matter, aimed at understanding the emergence of collective motion or other macroscopic transport processes in bacterial fluids, assume uncorrelated orientational noise, which is the direct consequence of the Poisson character of the R&T process [9].

The simple approach of introducing a Poisson distribution for the run times, although useful for simple qualitative interpretations, is not fully consistent with a growing number of measurements performed on the individual rotary motors [21–25] driving the helix-shaped flagella. For *E. coli*, the forward (run) motion is associated with the counterclockwise (CCW) rotation of the motors, and the tumbles take place when the motors rotate clockwise (CW). The CCW to CW transition is regulated by an internal biochemical process associated with the phosphorylation of the CheY protein.

In a seminal work, Korobkova *et al.* [21] brought evidence for a heavy tail distribution for the duration of CCW rotations. Importantly, this highlights possible coupling between the stochastic fluctuations in the chemotactic biochemical network and the emergent bacterial motility. Its consequences could affect the macroscopic organization of bacterial populations, chemotactic response to chemical heterogeneity, and genetic and epigenetic feedback of bacterial populations to environmental constraints.

Its potential importance in the context of active matter studies remains overlooked. For multiflagellated bacteria, the correspondence between switching statistics, motor synchronization, flagellar bundling and unbundling dynamics, and, finally, large-scale exploration properties remains unclear. Recently, indirect experimental evidence suggested that the macroscopic motility of free-swimming bacteria is sensitive to the stochasticity borne by the chemotactic biological circuit [26]. Here, we give direct evidence of this sensitivity.

Conceptually, our analysis starts from the extreme sensitivity of the rotational CCW \rightarrow CW switching to the abundance of the phosphorylated protein CheY-P in the cell. This picture induces a timescale separation, since, at short times, the alternation of CCW and CW rotations keeps a memory of a quasifixed level of CheY-P. This memory is erased at longer times, and we thus expect very different run times and motility features at the macroscopic level.

For the first time, we link the individual motor rotation statistics to the global motility features that we observe in a large number of 3D trajectories of wild-type *E. coli* bacteria. At short observation times, the time persistence of the swimming orientations displays an exponential decay as classically admitted, but with a large distribution of characteristic times within a population of monoclonal bacteria. However, when tracking the cells individually over several tenths of minutes, we identify for each

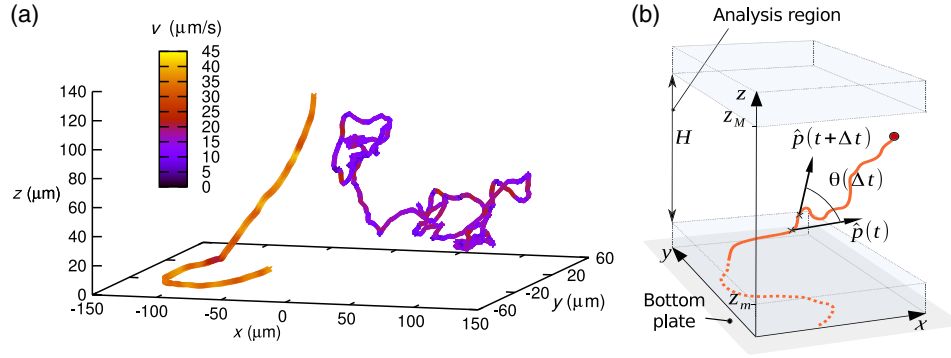
cell a large behavioral variability. The motility data are quantitatively analyzed through a simple model initially proposed by Tu and Grinstein [27] involving the fluctuations of CheY-P triggering the tumbling events. The model is here adapted to render the spatial exploration process. It now explains the occurrence of a large behavioral variability of swimming direction and also why, at short observation times, a large distribution of these is expected over a population. The central outcome of this model is that the persistence time durations naturally follow a log-normal distribution, instead of a standard Poisson distribution. Importantly, we identify a source of measurement bias introduced in most practical situations that is a consequence of such a large distribution of run times. Finally, we discuss the consequences of measuring averaged quantities over a population displaying a large distribution of motility features. This source of measurement bias is relevant in the general framework of experiments on statistical physics of active matter.

II. VARIABILITY OF BACTERIAL MOTILITY IN A POPULATION

To characterize the bacterial motility, we build an automated tracking device suited to follow fluorescent objects and record their 3D trajectories. A swimming bacterium is kept automatically in the center of the visualization field and at the focus of an inverted microscope by a visualization feedback loop acting horizontally on a mechanical stage and vertically on a piezo stage. The method is fully detailed in Ref. [28] by Darnige *et al.* (see also Sec. VI) and was recently used to investigate the swimming of bacteria in a Poiseuille flow [29].

We first monitor more than 100 swimming *E. coli* from different strains (see Sec. VI) in homogeneous diluted suspensions (concentration $\sim 10^5$ bact mL) confined between two horizontal glass slides, 250 μm apart. Figure 1(a) shows two typical trajectories from the same batch of monoclonal wild-type *E. coli*. We center our analysis on pieces of tracks exploring the bulk [Fig. 1(b)], i.e., in a measurement region located 10 μm above the surface and of maximum height $H = 130 \mu\text{m}$. For this series of experiments, the duration of a track is at minimum 8 s. We name these *experiments I*.

The bacterial velocities $\vec{V}(t)$ at each point of the trajectories are obtained by fitting the sequence of coordinates, along X , Y , and Z independently, over segments spanning 0.1 s, using a second-order polynomial. The first derivative of the polynomial evaluated at the center of the segment provides the velocity component. Figure 2 shows an example of a 3D trajectory and its velocity. Typically, the velocity curves for each track are irregular [Fig. 2(b)]. For a single track, the velocity distribution [Fig. 2(c)] shows a peak corresponding to the run phase and a low-velocity tail that might correspond to tumbling events. For the wild-type strain RP437 in a motility buffer, the average of the peak



F1:1 FIG. 1. Lagrangian 3D tracking of bacteria and analysis conditions. (a) RP437 wild-type *E. coli* displaying very different typical
F1:2 trajectories: persistent trajectory (bact 1: $\tau_p = 12$ s) and nonpersistent trajectory (bact 2: $\tau_p = 0.7$ s). (b) Sketch of the part of the track
F1:3 used for analysis and angles used for computing $C(\Delta t) = \langle \hat{p}(t) \cdot \hat{p}(t + \Delta t) \rangle = \langle \cos \theta(\Delta t) \rangle$, using a sliding window for an average on
F1:4 time t .

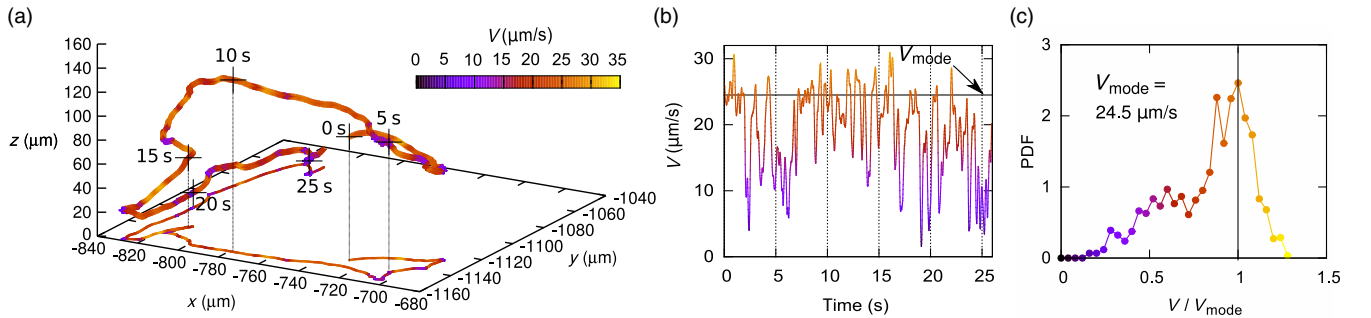
168 values for $V = |\vec{V}(t)|$ over the different tracks is $\langle V \rangle =$
169 $27 \pm 6 \mu\text{m/s}$.

170 Standard analysis to extract run-time distributions relies
171 on the identification of tumbling events, usually done by
172 detecting velocity drops and/or abrupt changes in the
173 swimming direction [2,6,30]. However, as shown in
174 Figs. 2(a) and 2(b), abrupt direction changes can take
175 place without a representative velocity decrease, and
176 velocity drops are sometimes not associated with reorien-
177 tation. This observation is consistent with results from
178 Refs. [30,31]. Moreover, by directly monitoring the flag-
179 ellar dynamics, Turner *et al.* [31] identify partial flagellar
180 debundling inducing weak velocity drops and directional
181 changes. We find that, without a direct observation of the
182 flagella, run-and-tumble detection requires the choice of
183 arbitrary criteria. We demonstrate this arbitrariness in the
184 Appendix A.

185 Here, in order to characterize the motility features, we do
186 not seek to explicitly identify the tumbling events. Instead,
187 we use the orientation correlation function $C(\Delta t)$ as a direct
188 measurement of the swimming direction persistence. The
189 director vectors pointing along the track are determined as
190 $\vec{p} = \vec{V}(t)/V(t)$ for each track. For each trajectory, we
191 compute $C(\Delta t) = \langle \hat{p}(t) \cdot \hat{p}(t + \Delta t) \rangle = \langle \cos[\theta(\Delta t)] \rangle$, where

θ is the angle between swimming directors separated by a
time lag Δt [Fig. 1(b)]. The brackets denote an average over
a time window sliding along the track. To ensure good
statistics, the maximum lag time Δt is chosen as one-tenth
of the total track duration. The orientation correlation
reflects the R&T statistics but advantageously does not
require an *ad hoc* criterion. In Fig. 3(a), 30 orientation
correlation functions obtained from separate tracks of
different bacteria (RP437 wild type in M9G) are displayed
as a function of Δt .

From the classical picture of an exponential distribution
of run times, the orientation correlation function is expected
to decay exponentially with a typical decay time of τ_p ,
defining the persistence time of the trajectory. For a
characteristic run time of $\bar{\tau}_{\text{run}} = 1$ s and a distribution of
reorientation angles of mean value $\theta_m = 51^\circ$ [1], one finds
 $\tau_p = (\bar{\tau}_{\text{run}}/1 - \langle \cos(\theta) \rangle) = 1.5$ s [32]. Recently, a slight
dependence of this angle on the swimming speed was
demonstrated [33] but is neglected in our study. Taking into
account rotational Brownian diffusion during the run phase
also leads to an exponential decaying correlation function
(see Appendix B). Its contribution represents a slight
modification to τ_p due to the much longer timescales of
Brownian diffusion. The predicted correlation function is



F2:1 FIG. 2. Details of a typical trajectory. (a) 3D trajectory and its projection on the x - y plane, (b) velocity vs time, and (c) velocity
F2:2 distribution. The marks every 5 s in the 3D track are references for comparison with (b).

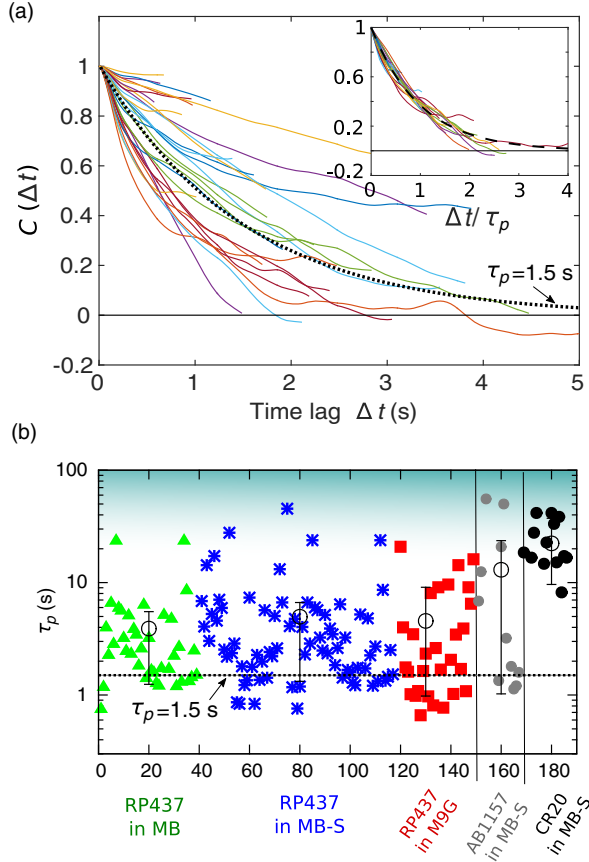


FIG. 3. Swimming orientation correlations. Experiments I. (a) Correlation function $C(\Delta t)$ obtained for 30 tracks of different RP437 bacteria in M9G, showing a large distribution of persistence times. The correlation functions are fitted with an exponential decay $\exp(-\tau/\tau_p)$ to extract the persistence times τ_p . The dotted line corresponds to $\tau_p = 1.5$ s as expected from Ref. [1]. Inset: Correlation functions as a function of Δt rescaled by τ_p . The dashed line is $\exp(-x)$. (b) Persistence times for individual bacteria of wild-type strains RP437 and AB1157 and smooth swimmer mutant CR20 in different media (MB, MB-S, and M9G). Circles and uncertainty bars correspond to the mean and 68% confidence intervals for each group. The blue background region designates the cutoff from Brownian diffusion. The dotted line corresponds to the expected $\tau_p = 1.5$ s also represented in (a). Uncertainty bars indicate the mean and confidence interval at 68%.

represented by the dotted line in Fig. 3(a). Strikingly, the experimental curves display a broad scattering, indicating a very large distribution of persistence times within this monoclonal population of bacteria.

By fitting the correlation functions with an exponential decay $\exp(-\tau/\tau_p)$, we determine the persistence times τ_p for each track. In Fig. 3(b), we display them on a logarithmic vertical axis for the strain RP437 in a motility buffer (MB) and a MB supplemented with serine (MB-S). In addition, persistence times obtained in a richer medium (M9G) and for a different wild-type strain AB1157 in

MB-S are shown. The results prove that the distribution of orientation persistence times for wild-type bacteria is very large. Within statistical errors, they are independent of the chemical environment (poor or rich), but they could depend on the strain, being larger in average for the 11 measurements performed on AB1157. For the very persistent tracks, the observed decorrelation remains weak over the accessible time lags. The obtained persistence times thus have a significant uncertainty, but we can be sure that their decorrelation time will be at least bigger than the time span of the track ($\tau_p > 8$ s). Finally, we consider the strain CR20, a smooth swimmer that tumbles only very rarely. In this case, the time distribution is gathered around the average $\tau_p = 25 \pm 10$ s, which is close to the Brownian rotational diffusion constant $\tau_p = \tau_B = 1/2D_r^B$, as expected. This value is, however, strongly dependent on the bacterial dimensions and aspect ratio [34,35]. A bacterium modeled as an ellipsoid of semiaxes $a = 4 \mu\text{m}$ and $b = c = 0.4 \mu\text{m}$ has a persistence time $\tau_p \sim 22$ s, while with $a = 6 \mu\text{m}$ it has a persistence time 3 times larger, $\tau_p \sim 66$ s [36]. Therefore, the wide distribution of persistence times for CR20 could arise from the bacterial size distribution. A possible origin of this dispersion on the measurement protocol is discussed in Sec. IV C.

III. VARIABILITY OF INDIVIDUAL BACTERIAL MOTILITY OVER TIME

The large diversity of trajectories here observed over short times in bacterial populations leads to the question of its origin. The diversity could arise from a phenotype multiplicity present in the monoclonal population [37,38], where each bacterium is characterized by a mean run time; alternatively, it could be due to temporal variability of the bacterial behavior, with mean run times varying over the course of time. To determine which scenario is taking place, we perform a second series of measurements, *experiments II*, where we follow individual bacteria over very long times (up to 20 min). In the new configuration, the top and bottom of the measurement chamber are within the observation range or the 3D tracker device. We follow individual bacteria as they alternate between the surfaces and the bulk, as sketched in Fig. 4(a). For the analysis, individual tracks are cut in pieces localized entirely in the bulk ($10 \mu\text{m}$ away from the walls). For each piece, we extract the persistence time from the correlation function. Finally, for each bacterium, we obtain a list of persistence times as a function of time. If the population displayed a large distribution of fixed run-times, one would expect for each bacterium a sequence of persistence times narrowly distributed around a characteristic value, but this value would be different for different bacteria. Figure 4(b) carries a very different message. For each of the tracks tested, the persistence times span a range of the same magnitude as

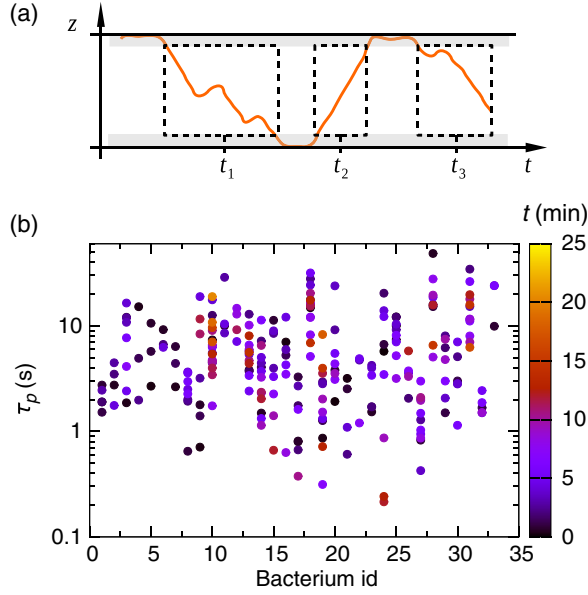


FIG. 4. Analysis for long tracks. Experiments II. (a) Sketch indicating the pieces of track from the same trajectory selected for demonstrating the behavioral variability. (b) τ_p for pieces of track from the same trajectories, for 33 different RP437 bacteria in MB-S. The color represents the starting time of the measurement. Each bacterium displays a large variation of persistence times.

for the whole population using shorter tracking times (see Fig. 3).

Previous studies based on 3D Eulerian tracking techniques [33,39], i.e., on a fixed reference frame, or even the Lagrangian tracking technique [6] were limited to short observation times and, consequently, were not able to catch such slow fluctuations of the run time. The fact that for a given bacterium the sequence of persistence times is largely distributed confirms the importance of behavioral variability in the motility process. However, due to tracking time limitations imposed by the bleaching of the fluorescent signal, we are not able to test precisely to what extent the behavioral variability contains features which could vary from one bacterium to the other, stemming from inherent phenotype variations, as identified, for example, by Dufour *et al.* [40].

IV. MOTILITY AND MOTOR ROTATION STATISTICS

The presence of a behavioral variability, as identified earlier, raises the question of its biochemical origins. Previous results point toward a definite influence of a stochastic process in the chemotactic sensory circuit. At the end of the biochemical cascade, there is a phosphorylation of a CheY protein (CheY-P) promoting a switch in the motor rotation from the CCW state (run phase) to the CW state (tumbling phase). The most accepted picture rendering the $\text{CCW} \rightleftharpoons \text{CW}$ transition is a two-state model initially proposed by Khan and Macnab [41], which considers the

switching of the rotation direction $\text{CCW} \rightarrow \text{CW}$ (equivalently, $\text{CW} \rightarrow \text{CCW}$) as an activated process regulated by the presence of CheY-P. The double-well Gibbs free energy associated with the transition $\text{CCW} \rightleftharpoons \text{CW}$ depends in a very sensible way on the CheY-P ($[Y]$) concentration values near the motor, as shown by Cluzel, Surette, and Leibler [42]. This strong sensitivity leads naturally to behavioral variations, as slow fluctuations around the mean value can change the motility features from preferentially tumbling (high CheY-P) to preferentially running (low CheY-P). It also means that at short times the CheY-P level does not change significantly and motility features remain constant. Therefore, at a given moment, the motility features should be largely distributed in a population of bacteria bearing different CheY-P concentrations. This large distribution is in essence what is observed in our experiments in Figs. 3 and 4.

A. Quantitative description of the behavioral variability model

To rationalize and quantify our experimental findings, we adapt the simple but enlightening physical model proposed by Tu and Grinstein [27]. The behavioral variability (BV) model we present here quantifies the role of fluctuations of the phosphorylated protein CheY-P in the regulation of the motor-switching statistics. The key idea is that the observed typical switching time at a given moment depends on the instantaneous CheY-P concentration $[Y](t)$. Then, considering concentration fluctuations around a mean value ($\delta Y(t) = [Y](t) - [Y_0]$), one obtains a two-state model with a time-varying barrier describing the $\text{CCW} \rightarrow \text{CW}$ switching process. Tu and Grinstein model the δY fluctuations as an Ornstein-Uhlenbeck process with a memory (relaxation) time T_Y , hence yielding a Gaussian distribution for δY values. Note that T_Y is considered to be larger than typical motor-switching times [see Fig. 5(a) for the relevant timescales].

For small fluctuations of concentration, the average switching time can be written as

$$\tau_s = \tau_0 e^{-\Delta_n \delta X}. \quad (1)$$

Here, δX corresponds to the fluctuations in concentration normalized by the δY standard deviation σ_Y ; τ_0 is a typical switching time corresponding to the mean concentration $[Y_0]$ and $\Delta_n = \alpha(\sigma_Y/Y_0)$. The parameter α is positive [42] and measures the sensitivity of the switch to variations in $[Y]$, which means that higher concentrations of CheY-P lead to shorter run times. Note that, in principle, the two switching times describing $\text{CCW} \rightarrow \text{CW}$ (run times) or $\text{CW} \rightarrow \text{CCW}$ (tumbling times) could be modeled with corresponding parameters τ_0 and Δ_n . However, as the results from Korobkova *et al.* [21] show that, in contrast with run times, the distribution of tumble times is exponential, meaning that the equivalent of Δ_n for tumbles is

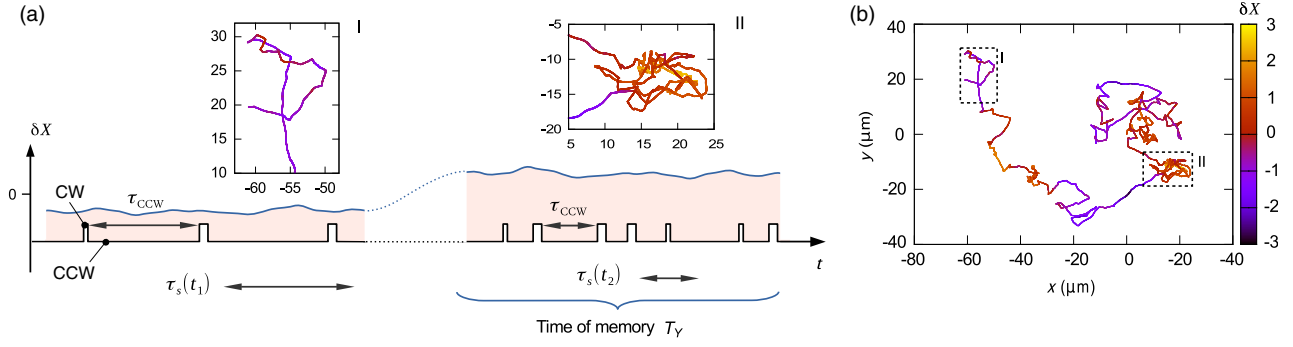


FIG. 5. Heuristic view of the behavioral variability model. (a) Timescales of the tumbling process and the CheY-P concentration governing them. The switching time τ_s represents the local average of the stochastic run times τ_{CCW} . The switching time τ_s stays relatively constant during the memory time T_Y and evolves as a function of the normalized CheY-P concentration: $\delta X = ([Y] - [Y_0]) / \sigma_Y$. (b) 2D projection of the simulated 3D trajectory where the δX fluctuations drive the tumbling process. The insets correspond to different levels of $[\delta X]$: Inset I depicts a low CheY-P level, and inset II depicts a high CheY-P level.

small. Hence, we consider the tumbling times as a Poissonian process, well described by a single timescale.

Let us first consider the $CCW \rightarrow CW$ switching time distributions. Each observed time belongs to a Poisson distribution with a typical time τ_s set by the current CheY-P concentration $[Y](t)$ [see Eq. (1)]. As a consequence, the observed switching statistics for an individual bacterium, when observed over a time interval shorter than the memory time, should approximately appear as an effective Poisson process, which is indeed the case, as shown from the collapse of the rescaled orientation correlation functions onto a single exponential decay shown in Fig. 3(a). The model provides a second important outcome. A random choice of a bacterium in a population is like a random choice of δX , hence defining a typical switching time τ_s for this bacterium. A Gaussian distribution for δX , as assumed by the BV model, leads to a Gaussian distribution of $\ln(\tau_s)$ characterized by an average $\ln(\tau_0)$ and a standard deviation $\sigma_{\ln \tau_p} = \Delta_n$, yielding naturally a large log-normal distribution of τ_s provided the switch

sensitivity α is large. Note that the power law distribution discussed by Tu and Grinstein [27] is obtained in the limit of very large Δ_n and not in contradiction with the above statement. As τ_s and τ_p are proportional, the distribution of $\ln(\tau_p)$ should also be Gaussian.

To illustrate this idea, a very long 3D trajectory is synthesized numerically using the switching statistics from the BV model. Figure 5(b) shows a 2D projection (see Sec. VI for technical details and Sec. IV C for the parameter values). The simulated trajectory contains very persistent (inset I) and very nonpersistent (inset II) parts. The colors represent the local values of δX illustrating the direct influence of the slow variations of CheY-P concentration on the bacterial motility, hence explaining the observed behavioral variability.

B. Memory time

The evolution of persistence times τ_p along individual trajectories displays large variations. It is shown in Fig. 6(a)

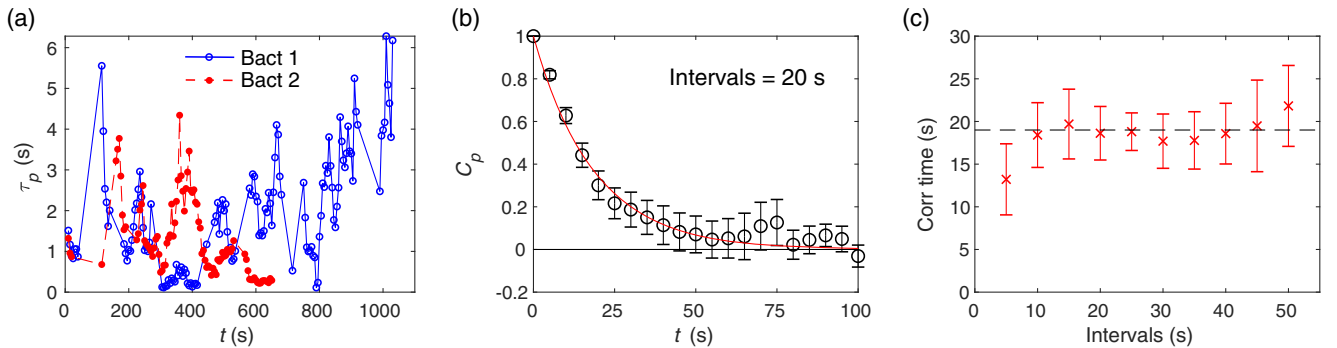


FIG. 6. Determination of the memory time from experiments II. (a) Persistence times τ_p computed over pieces of span 20 s and shifted 5 s for two different bacteria. Gaps larger than 5 s between consecutive points correspond to lapses in which the bacteria are swimming close to surfaces. (b) Persistence times self-correlation function C_p using pieces of 20 s. Points represent the average over the ensemble of bacteria. (c) Correlation time of the persistence times as a function of the lengths of the pieces. We extract the memory time to be $T_Y = 19.0 \pm 1.3$ s.

for the case of two different bacteria continuously tracked for 11 and 17 min. The values of τ_p for each track are extracted from intervals of span 20 s shifted 5 s along the trajectory. Gaps larger than 5 s between consecutive points correspond to lapses in which the bacterium is swimming close to a surface. Analyzing, for example, the bacterium of the blue longer trajectory, at time 300 s (5 min) it displays a persistence time close to 0.1 s, in contrast with a persistence time close to 5 s around time 1000 s (~ 17 min). This temporal variation of τ_p is considered in the framework of the BV model. The memory time T_Y is then a central parameter of the BV model, as it provides a natural separation between short-time measurements and long-time measurements. Therefore, for a correct statistical interpretation of the results, τ_p values must be extracted from pieces of tracks not longer than the memory time T_Y .

We estimate the memory time T_Y from the long-time tracking data in experiments II, using the following procedure. For each bacterial trajectory, we compute a sequence of τ_p using intervals of a specific duration. For each sequence of τ_p , we compute the self-correlation function of persistence times, $C_p(t) = \langle \ln \tau_p(t + t') \ln \tau_p(t') \rangle - \langle \ln \tau_p \rangle^2$, where the average is done over t' . The average of C_p over the ensemble of trajectories is fitted with an exponential, giving the correlation time [Fig. 6(b)]. With this procedure, we investigate different lengths of intervals [Fig. 6(c)], finding that the correlation times grow with the duration of the interval until saturation at the value of the memory time $T_Y \approx 19.0 \pm 1.3$ s.

C. Comparison with the model

The BV model depends on several parameters: the memory time T_Y , the mean switch time and sensitivity τ_0 and Δ_n , respectively, the rotational diffusion coefficient D_r^B , and the dimensionless rotational diffusion coefficient \tilde{D}_r^{eff} used to model the reorientation during tumble (see Sec. VI for details). We determine T_Y from the experiments, while the rest of the parameters are fitted using the following protocol. A long simulated trajectory is generated and cut in pieces of duration 20 s, similar to the analysis of the experimental tracks, and the persistence time is computed for each piece. We look for the values of the parameters that best agree with the experimental values of the first four moments of the distribution of $\ln \tau_p$. The result is $\tau_0 = 1.53$ s, $\Delta_n = 1.62$, $D_r^B = 0.025$ s $^{-1}$, and $\tilde{D}_r^{\text{eff}} = 3.86$. Note that the velocity does not appear in the fit, because we compare simulations and experiments using the persistence times, which depend only on the orientations.

Figure 7(a) compares the experimental distribution of $\ln \tau_p$ with the results from simulations using the optimal parameters. The agreement is very good, with two features that need discussion. First, in agreement with the BV model, the distributions are not exactly Gaussian but

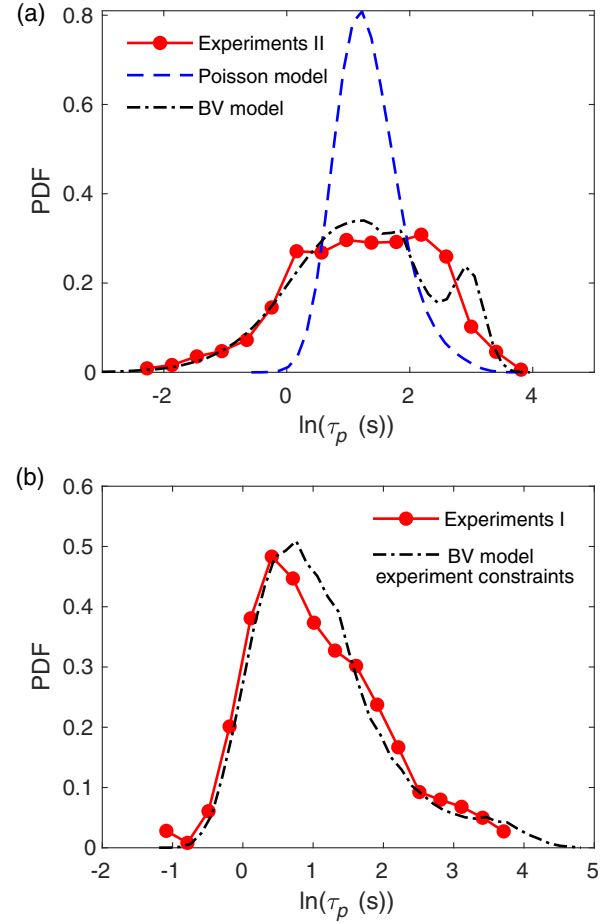


FIG. 7. Probability density function of the logarithm of persistence times ($\ln \tau_p$). (a) The values τ_p are extracted from pieces of track from experiments II that last 20 s. Simulations using two different models are shown: The Poisson model does not reproduce the experiments, while the BV model reproduces the main features. (b) The experimental distribution corresponds to the combined RP437 bacteria in all media, from Fig. 3 in experiments I. “BV model experiment constraints” are determined from the same simulations of “BV model” but analyzing pieces of trajectories that follow the experimental constraints in this case. It reproduces experiments I without additional free parameters.

present a negative excess kurtosis. With 63% probability, the switch times are in the range $[\tau_0 e^{-\Delta_n}, \tau_0 e^{\Delta_n}] = [0.30 \text{ s}, 7.7 \text{ s}]$. Hence, there is no complete separation of timescale with T_Y . As a consequence, in each piece, δX is not constant, and the measured and simulation distributions result from the mixture of different values of τ_s . Note that shorter pieces would imply too few tumble events and would make it unreliable to use the orientation correlation function. A perfect log-normal distribution could be observed if there was a good separation of timescales, allowing a choice of intervals such that $\tau_0 \ll T_{\text{interval}} \ll T_Y$.

The second feature is the small peak at $\ln \tau_p \approx 3$ in the simulations. This peak corresponds to pieces of the

trajectory where no single tumble took place. The change in orientation is due only to rotational diffusion during a run. Because $\tau_B = 1/2D_r^B \approx 20$ s is similar to T_Y , no complete reorientation occurs in the interval, resulting in a distribution of τ_p for nontumbling swimmers. In fact, the persistence times for the nontumbling bacteria [strain CR20 from experiments I, Fig. 3(b)] coincide with this peak. This feature should also be present in experiments, but, as discussed in Sec. II, D_r^B depends strongly on the bacterial dimensions, which vary within the population. This dispersion of rotational diffusion and other imperfections blur this peak in the experiments, contrary to the simulations, where all swimmers are identical. Note that, despite of the diversity, the fitted value of D_r^B matches closely the prediction made in Sec. II for ellipsoidal swimmers.

Since the pieces of trajectories are of a finite length, the orientation correlation function is not perfectly sampled, and, even for a constant switch time τ_s , the persistence times τ_p obtained from an exponential fit would present some dispersion. To test whether the observed dispersion is due only to the data analysis protocol, we perform simulations with a Poisson model. For this test, we look for the best parameters to reproduce the first fourth moments of the distribution of $\ln \tau_p$, setting $\Delta_n = 0$. The result is $\tau_0 = 1.18$ s, $D_r^B = 0.026$ s⁻¹, and $\tilde{D}_r^{\text{eff}} = 1.61$. Figure 7(a) presents the resulting distribution, which is far from the experimental one. We conclude that a Poisson process cannot explain the broad distribution of persistence times observed experimentally.

Finally, for consistency, we return to the persistence times obtained in Fig. 3 from experiments I. In this experimental protocol, the trajectories are selected within a certain height (10–140 μ m from the surface) and longer than 8 s. The corresponding experimental distribution of $\ln(\tau_p)$ for RP437 bacteria in all media [Fig. 7(b)] displays a clear positive skewness, which differs strikingly from the experimental measures of Fig. 7(a), done using the same bacterial strain and confinement and a similar chemical environment. This difference originates from a measurement bias built into the analysis of Fig. 7(b) (and Fig. 3). The bias is a consequence of a preferential selection of long trajectories staying essentially in the x - y plane, with limited bounds in the vertical direction. The skewness is enhanced by the broad distribution of run times, since very persistent swimmers will likely quit the measurement region in a very short time, hence privileging small persistence times. The curve “BV model” represents the distribution of persistence times from simulations of the BV model that fit the experiments in Fig. 7(a) (experiments I). When this same simulation is analyzed by taking pieces following strictly the experimental constraints, on both duration and vertical spatial exploration, the resulting distribution (“BV model experiment constraints”) compares very well and noticeably without any additional fitting parameter, to the experimental curve in Fig. 7(b) (experiments II).

V. CONCLUSIONS

We have shown that the 3D spatial exploration of an adapted *E. coli* reflects a behavioral variability that we associate with intrinsic noise in the chemotaxis pathway controlling the run-and-tumble sequence. Our results for free-swimming bacteria are consistent with models describing motor-switching dynamics based on tethered cell measurements. We identified a large log-normal distribution of persistent times stemming from the slow fluctuations of an internal variable accounting for the CheY-P concentration near the motors. In the context of many recent works on statistical physics of active matter, we suggest that this large variability should be included in the description of bacterial fluids. This variability is expected to influence the computation of averaged quantities like diffusivity, viscosity, or any constitutive relations of macroscopic transport processes.

The broad distribution of run times is likely to introduce measurement biases in practical situations. Here, we reduce the bias by taking pieces of trajectories of equal length, not larger than the memory time. Mixing trajectories of different lengths can result in highly distorted distributions.

The large distribution of motility features has been related to the time bacteria spend close to surfaces. As an example, the existence of a large distribution of motor-switching statistics was found crucial to understand large-scale upstream bacterial contamination of narrow channels [26], where substantial transport occurs along surfaces [43–46].

We expect the chemotactic drift to be sensitive to the distribution of CheY-P concentrations, since a nonlocal spatiotemporal coupling will take place between chemical gradients and bacterial concentration. This sensitivity should be taken into account in future motility modeling. Finally, these findings may also impact quantitative modeling on how bacterial populations react to environmental changes, colonize space, swarm in a biofilm [47], or interact with other communities.

VI. MATERIALS AND METHODS

A. Bacterial strains and culture

We use the wild-type strains RP437 and AB1157 and a smooth swimmer mutant strain CR20 (Δ CheY) expressing yellow fluorescent protein (YFP) from a plasmid. Bacteria are grown overnight at 30 °C in M9G medium [M9 minimal medium supplemented with glucose (4 g/L), casamino acids (1 g/L), MgSO₄ (2 mM), and CaCl₂ (0.1 mM)] plus the corresponding antibiotics, up to optical density = 0.5 at 595 nm. Cells are then washed 3 times by centrifugation at 2000 g for 5 min and suspended in a motility buffer (10 mM potassium phosphate buffer pH ~ 7.0, 0.1 mM EDTA, 1 μ M L-methionine, and 10 mM sodium L-lactate), supplemented with polyvinylpyrrolidone (PVP-360 kDa 0.002%) and, when indicated, with L-Serine (0.04 g/mL).

B. The 3D Lagrangian tracker

We develop a device for keeping individual microscopic objects—as swimming bacteria—in focus, as they move in microfluidic chambers [28]. The system is based on real-time image processing, determining the displacement of the stage to keep the chosen object at a fixed position in the observation frame. The z displacement of the stage is based on the refocusing of the fluorescent object that keeps the moving object in focus. The algorithm for z determination is designed for not being affected by photobleaching.

The instrument is mounted on an epifluorescent inverted microscope (Zeiss-Observer, Z1) with a high magnification objective ($100\times/0.9$ DIC Zeiss EC Epiplan-Neofluar), an x - y mechanically controllable stage with a z piezomover from Applied Scientific Instrumentation (ms-2000-flat-top-xyz), and a digital camera ANDOR iXon 897 EMCCD. The device works nominally at 30 frames per second on a 512×512 pix² matrix, but a faster tracking speed of 80 Hz can be achieved by reducing the spatial resolution to 128×128 pix². It provides images of the object and its track coordinates with respect to the microfluidic device.

The tracking limitations come essentially from the z exploration range, restricted by the working distance of $150\text{ }\mu\text{m}$ of the objective. In the x - y plane, the spatial limitations are virtually nonexistent, since the stage displacement can be as long as 15 cm, which is much bigger than the typical sizes of the sample (a few millimeters). Details of the apparatus are given in Ref. [28], as well as an exhaustive explanation of a method for correcting the mechanical backlash typically affecting these systems and a discussion of the device's performance and limitations.

C. Experimental geometries and bacteria tracking

We monitor hundreds of single *E. coli* in a drop of a diluted homogeneous suspension (concentration $\sim 10^5$ bact/mL) squeezed between two horizontal glass slides. The drop has typically a diameter of 1 mm. The gap between the two glass plates is $250\text{ }\mu\text{m}$. For experiments I, displayed in Fig. 3, only pieces of 3D trajectories remaining between the vertical bounds $z_m = 10\text{ }\mu\text{m}$ from the bottom surface and $z_M = 140\text{ }\mu\text{m}$, the highest possible height, and lasting more than 8 s are taken into account. For the set of very long tracks in Fig. 4, experiments II, the gap between the glass plates is also $250\text{ }\mu\text{m}$, but the whole trajectories are captured, as they alternate between the bottom and top. For the analysis, only pieces farther than $10\text{ }\mu\text{m}$ from the surfaces are taken into account.

The velocities are determined from second-order Savitzky-Golay filtering of the coordinates over 0.1 s, resulting in uncertainties close to 5% [36]. For each track, the velocity distribution shows a peak corresponding to the mean run velocity and a low-velocity tail corresponding to the contribution of sudden velocity drops (Fig. 2). Peak

velocities are on average $\langle V \rangle = 27 \pm 6\text{ }\mu\text{m/s}$. To compute the correlation function $C(\Delta t)$, the average is made over time, and the lag time is offset by 0.2 s to avoid the short-time decorrelation due to wobbling [36,48]. The correlation function is then normalized by its value at 0.2 s to yield 1 at the lag time origin.

D. Track simulations using the BV model

Swimmers are described by their position \vec{r} , orientation \hat{p} , and the instantaneous value of the normalized fluctuations of the CheY-P concentration $\delta X = ([Y] - [Y_0])/\sigma_Y$. During the run phase, they obey the equations

$$\dot{\vec{r}} = V\hat{p}, \quad (2)$$

$$\dot{\hat{p}} = \sqrt{D_r^B}(I - \hat{p}\hat{p})\vec{\eta}(t), \quad (3)$$

$$\dot{\delta X} = -\delta X/T_Y + \sqrt{2/T_Y}\xi(t), \quad (4)$$

where V is the swim velocity, D_r^B is the rotational diffusion coefficient, T_Y is the memory time, $(I - \hat{p}\hat{p})$ is a projector orthogonal to \hat{p} , ξ is a white noise of zero mean and correlation $\langle \xi(t)\xi(s) \rangle = \delta(t-s)$, and $\vec{\eta}$ is a white noise vector of zero mean, where the components have correlations $\langle \eta_i(t)\eta_k(s) \rangle = \delta_{ik}\delta(t-s)$.

The BV model yields a relation between the characteristic switching time for the transition CCW \rightarrow CW (run to tumble) and the CheY-P concentration. As a simplification, we assume that, due to the small cellular dimensions, all six flagella operate at the same CheY-P concentration and that the reverse of the rotation direction of a single flagellum is enough to trigger a tumble. Hence, the probability to tumble in Δt would be $6\Delta t/\tau_s$. To simplify notation, we absorb the factor 6 into τ_0 , resulting in a tumble probability $\Delta t e^{\Delta_n \delta X}/\tau_0$.

The BV model predicts that the characteristic switching time for the transition CCW \rightarrow CW (tumble to run) is also given from an activated process. But, as the corresponding value of Δ_n is small, the tumble duration is given by a Poisson process with characteristic time τ_1 . In addition, the reorientation dynamics during a tumble needs to be modeled. *A priori*, the link between motor switch and tumble is far from being trivial, as, in principle, one needs to account for the hydrodynamically complex bundling and unbundling process of the multiflagellated *E. coli* bacteria [49,50]. Here, we rather follow a simple effective approach inspired by Saragosti, Silberzan, and Buguin [51]. We model the reorientation dynamics during tumbling as an effective rotational diffusion process with a coefficient D_r^{eff} . Defining the dimensionless combination $\tilde{D}_r^{\text{eff}} = D_r^{\text{eff}}\tau_1$, the dimensionless tumble durations are sorted from an exponential distribution with a typical time equal to one, and, during a tumble, the dynamics is

$$\dot{\vec{r}} = 0, \quad (5)$$

$$\dot{\hat{p}} = \sqrt{\tilde{D}_r^{\text{eff}}} (I - \hat{p} \hat{p}) \vec{\eta}(t), \quad (6)$$

$$\delta \dot{X} = 0. \quad (7)$$

After the tumble phase, a new run phase starts.

ACKNOWLEDGMENTS

The authors thank Dr. Reinaldo García García for useful discussions, Professor Axel Bugin for bacterial strains, and Professor Igor Aranson for comments on the manuscript. This work was supported by the ANR grant “BacFlow” ANR-15-CE30-0013 and the Franco-Chilean Ecosud Collaborative Program C16E03. N.F.-M. thanks the Pierre-Gilles de Gennes Foundation for financial support. A.L. and N.F.-M. acknowledge support from the ERC Consolidator Grant PaDyFlow under Grant Agreement No. 682367. R.S. acknowledges the Fondecyt Grant No. 1180791 and Millennium Nucleus Physics of Active Matter of the Millennium Scientific Initiative of the Ministry of Economy, Development and Tourism (Chile). V.A.M. was funded by ERC AdG 340877 (PHYSAPS) and Joliot-Curie Chair from ESPCI. E.C. and R.S. thank the support from Laboratoire International Associé “Matière: Structure et dynamique” (LIA-MSD), CNRS-France. This work has received the support of Institut Pierre-Gilles de Gennes (Équipement d’Excellence, “Investissements d’avenir,” Program No. ANR-10-EQPX-34).

APPENDIX A: RUN DURATION ANALYSIS

Figure 8 displays a series of analyses on a single trajectory, evidencing that tumble detection is criterion dependent. Here, we set a threshold velocity, cutoff V , and identify as runs all the continuous parts of the track where the bacterial velocity is above the prescribed threshold. The plot demonstrates that, by changing the cutoff value for the velocity, we can obtain average run times of duration 1–8 s.

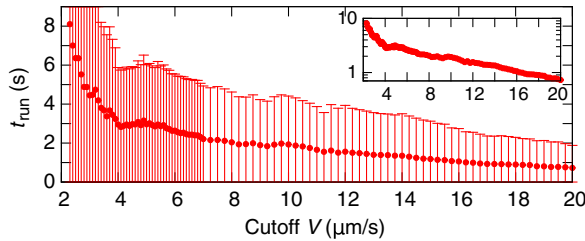


FIG. 8. Average run duration as a function of the threshold in bacterial velocity (cutoff V). The runs are identified as continuous parts of the track where $V > \text{cutoff } V$. The plot demonstrates that an arbitrary choice of velocity drops along the trajectory leads to arbitrary run-time duration. Inset: Log-normal plot.

APPENDIX B: PERSISTENCE CORRELATION FUNCTION

The orientation correlation function is defined as

$$C(\tau) = \langle \hat{p}(t) \cdot \hat{p}(t + \tau) \rangle = \langle \cos[\theta(\tau)] \rangle, \quad (\text{B1})$$

where \hat{p} is the director vector and the average is done over time t .

To compute the correlation function, we use a kinetic theory approach. The object under study is the distribution function $f(\hat{p}, t)$, which gives the probability that a bacterium has an orientation \hat{p} at time t . In this context, the correlation function is obtained assuming that the initial condition at $t = 0$ is with the bacterium pointing in a specific direction, say, \hat{p}_0 . Hence, we have to compute $C(\tau) = \langle \hat{p}(\tau) \cdot \hat{p}_0 \rangle$, where now the average is over the distribution function. At the end, another average, over \hat{p}_0 , should be done. In practice, this last average is unnecessary by the isotropy of space, because the first average gives already a value independent of \hat{p}_0 .

The distribution function obeys the kinetic equation [52,53]

$$\frac{\partial f}{\partial t} = -L f, \quad (\text{B2})$$

with

$$f(\hat{p}, 0) = \delta(\hat{p} - \hat{p}_0) \quad (\text{B3})$$

and L the evolution operator. Two models must be considered. In the case of Brownian rotational diffusivity,

$$L f = -D_r^B \nabla_{\hat{p}}^2 f, \quad (\text{B4})$$

where D_r^B is the rotational diffusion coefficient and $\nabla_{\hat{p}}^2$ is the angular part of the Laplacian. In the case of tumbling with a characteristic switch time τ_s ,

$$L f = \frac{1}{\tau_s} \left[f - \int d\hat{p}' W(\hat{p}', \hat{p}) f(\hat{p}') \right]. \quad (\text{B5})$$

The kernel $W(\hat{p}', \hat{p})$ gives the probability that for a swimmer with director \hat{p}' , after tumbling, the new director is \hat{p} . It is normalized to $\int d\hat{p} W(\hat{p}', \hat{p}) = 1$, indicating that some director \hat{p} must be chosen. If the space is isotropic, the kernel depends only on the relative angle between the directors, that is, $W(\hat{p}', \hat{p}) = w(\hat{p}' \cdot \hat{p})$. Finally, if tumbling and diffusion are present, the operator is just the sum of both.

If the space is isotropic, the evolution operator is also isotropic, which in this case implies that it commutes with the angular Laplacian $\nabla_{\hat{p}}^2$. Therefore, both operators share eigenfunctions, which are the spherical harmonics $Y_{lm}(\hat{p})$. Then, there are eigenvalues λ_l ,

$$LY_{lm} = \lambda_l Y_{lm}, \quad (\text{B6})$$

that, by isotropy, do not depend on the second index m . For the diffusion case, the eigenvalues are known exactly, while for tumbling they are proportional to $1/\tau_s$ and depend on the kernel model. In summary,

$$\lambda_l = D_r^B l(l+1) + 1/(a_l \tau_s), \quad (\text{B7})$$

where a_l are dimensionless parameters of the order of 1 that depend on the kernel w .

Using the basis of the spherical harmonics, the solution of the kinetic equation (B2) is

$$f(\hat{p}, t) = \sum_{lm} f_{lm}(0) Y_{lm}(\hat{p}) e^{-\lambda_l t}, \quad (\text{B8})$$

where $f_{lm}(0)$ depend on the initial condition (B3).

Now, the correlation function is

$$C(t) = \langle \hat{p}(t) \cdot \hat{p}_0 \rangle \quad (\text{B9})$$

$$= \int d\hat{p} \hat{p}_0 \cdot \hat{p} f(\hat{p}, t) \quad (\text{B10})$$

$$= \sum_{lm} f_{lm}(0) e^{-\lambda_l t} \hat{p}_0 \cdot \int d\hat{p} \hat{p} Y_{lm}(\hat{p}). \quad (\text{B11})$$

Using that \hat{p} can be written as a linear combination of Y_{lm} , with $m = 0, \pm 1$ and the orthogonality of the spherical harmonics, it is obtained that the integral is not vanishing only for $l = 1$. Combining factors, one obtains

$$C(t) = C_0 e^{-t/\tau_p}, \quad (\text{B12})$$

where

$$\tau_p = \frac{a_1 \tau_s}{1 + a_1 \tau_s / \tau_B} \quad (\text{B13})$$

and $\tau_B = 1/(2D_r^B)$ is the Brownian decorrelation time.

In the classical picture, where all bacteria have a single value for τ_s , the decorrelation time τ_p is single valued also. When τ_s is broadly distributed, the decorrelation time τ_p also follows a broad distribution for $\tau_p \ll \tau_B$, and it is bounded from above by τ_B . Finally, in the description of Berg and Brown [1], the tumble angles are distributed with a peak at 63° . In this case, $a_1 = 1/(1 - \langle \cos \theta \rangle)$ [51].

- [3] J. Saragosti, V. Calvez, N. Bournaveas, B. Perthame, A. Buguin, and P. Silberzan, *Directional Persistence of Chemotactic Bacteria in a Traveling Concentration Wave*, *Proc. Natl. Acad. Sci. U.S.A.* **108**, 16235 (2011).
- [4] U. Alon, L. Camarena, M. G. Surette, B. A. y Arcas, Y. Liu, S. Leibler, and J. B. Stock, *Response Regulator Output in Bacterial Chemotaxis*, *EMBO J.* **17**, 4238 (1998).
- [5] H. C. Berg, *Random Walks in Biology* (Princeton University, Princeton, NJ, 1993).
- [6] Z. Qu, F. Z. Temel, R. Henderikx, and K. S. Breuer, *Changes in the Flagellar Bundling Time Account for Variations in Swimming Behavior of Flagellated Bacteria in Viscous Media*, *Proc. Natl. Acad. Sci. U.S.A.* **115**, 1707 (2018).
- [7] X.-L. Wu and A. Libchaber, *Particle Diffusion in a Quasi-Two-Dimensional Bacterial Bath*, *Phys. Rev. Lett.* **84**, 3017 (2000).
- [8] D. Saintillan and M. J. Shelley, *Active Suspensions and Their Nonlinear Models*, *C.R. Phys.* **14**, 497 (2013).
- [9] M. C. Marchetti, J. F. Joanny, S. Ramaswamy, T. B. Liverpool, J. Prost, M. Rao, and R. A. Simha, *Hydrodynamics of Soft Active Matter*, *Rev. Mod. Phys.* **85**, 1143 (2013).
- [10] A. P. Solon, Y. Fily, A. Baskaran, M. E. Cates, Y. Kafri, M. Kardar, and J. Tailleur, *Pressure Is Not a State Function for Generic Active Fluids*, *Nat. Phys.* **11**, 673 (2015).
- [11] J. Schwarz-Linek, J. Arlt, A. Jepsen, A. Dawson, T. Vissers, D. Miroli, T. Pilizota, V. A. Martinez, and W. C. K. Poon, *Escherichia coli as a Model Active Colloid: A Practical Introduction*, *Colloids Surf. B* **137**, 2 (2016).
- [12] P. Galajda, J. Keymer, P. Chaikin, and R. Austin, *A Wall of Funnel Concentrates Swimming Bacteria*, *J. Bacteriol.* **189**, 8704 (2007).
- [13] C. Dombrowski, L. Cisneros, S. Chatkaew, R. E. Goldstein, and J. O. Kessler, *Self-Concentration and Large-Scale Coherence in Bacterial Dynamics*, *Phys. Rev. Lett.* **93**, 098103 (2004).
- [14] A. Sokolov and I. S. Aranson, *Reduction of Viscosity in Suspension of Swimming Bacteria*, *Phys. Rev. Lett.* **103**, 148101 (2009).
- [15] J. Gachelin, G. Miño, H. Berthet, A. Lindner, A. Rousselet, and E. Clément, *Non-Newtonian Viscosity of Escherichia coli Suspensions*, *Phys. Rev. Lett.* **110**, 268103 (2013).
- [16] H. M. López, J. Gachelin, C. Douarche, H. Auradou, and E. Clément, *Turning Bacteria Suspensions into Superfluids*, *Phys. Rev. Lett.* **115**, 028301 (2015).
- [17] A. Sokolov, M. M. Apodaca, B. A. Grzybowski, and I. S. Aranson, *Swimming Bacteria Power Microscopic Gears*, *Proc. Natl. Acad. Sci. U.S.A.* **107**, 969 (2010).
- [18] R. Di Leonardo, L. Angelani, D. DellArciprete, G. Ruocco, V. Iebba, S. Schippa, M. P. Conte, F. Mecarini, F. De Angelis, and E. Di Fabrizio, *Bacterial Ratchet Motors*, *Proc. Natl. Acad. Sci. U.S.A.* **107**, 9541 (2010).
- [19] H. Wioand, F. G. Woodhouse, J. Dunkel, J. O. Kessler, and R. E. Goldstein, *Confinement Stabilizes a Bacterial Suspension into a Spiral Vortex*, *Phys. Rev. Lett.* **110**, 268102 (2013).
- [20] A. Kaiser, A. Peshkov, A. Sokolov, B. ten Hagen, H. Löwen, and I. S. Aranson, *Transport Powered by Bacterial Turbulence*, *Phys. Rev. Lett.* **112**, 158101 (2014).

[1] H. C. Berg and D. A. Brown, *Chemotaxis in Escherichia coli Analysed by Three-Dimensional Tracking*, *Nature (London)* **239**, 500 (1972).

[2] H. C. Berg, *E. coli in Motion* (Springer, New York, 2004).

- [21] E. Korobkova, T. Emonet, J. M. G. Vilar, T. S. Shimizu, and P. Cluzel, *From Molecular Noise to Behavioural Variability in a Single Bacterium*, *Nature (London)* **428**, 574 (2004).
- [22] E. A. Korobkova, T. Emonet, H. Park, and P. Cluzel, *Hidden Stochastic Nature of a Single Bacterial Motor*, *Phys. Rev. Lett.* **96**, 058105 (2006).
- [23] T. Emonet and P. Cluzel, *Relationship between Cellular Response and Behavioral Variability in Bacterial Chemotaxis*, *Proc. Natl. Acad. Sci. U.S.A.* **105**, 3304 (2008).
- [24] T. L. Min, P. J. Mears, L. M. Chubiz, C. V. Rao, I. Golding, and Y. R. Chemla, *High-Resolution, Long-Term Characterization of Bacterial Motility Using Optical Tweezers*, *Nat. Methods* **6**, 831 (2009).
- [25] F. Wang, H. Shi, R. He, R. Wang, R. Zhang, and J. Yuan, *Non-equilibrium Effect in the Allosteric Regulation of the Bacterial Flagellar Switch*, *Nat. Phys.* **13**, 710 (2017).
- [26] N. Figueroa-Morales, A. Rivera, R. Soto, A. Lindner, E. Altshuler, and E. Clément, *E. coli “Super-Contaminates” Narrow Ducts Fostered by Broad Run-Time Distribution*, *Sci. Adv.* **6**, eaay0155 (2020), <https://advances.sciencemag.org/content/6/11/eaay0155>.
- [27] Y. Tu and G. Grinstein, *How White Noise Generates Power-Law Switching in Bacterial Flagellar Motors*, *Phys. Rev. Lett.* **94**, 208101 (2005).
- [28] T. Darnige, N. Figueroa-Morales, P. Bohec, A. Lindner, and E. Clément, *Lagrangian 3D Tracking of Fluorescent Microscopic Objects in Motion*, *Rev. Sci. Instrum.* **88**, 055106 (2017).
- [29] G. Junot, N. Figueroa-Morales, T. Darnige, A. Lindner, R. Soto, H. Auradou, and E. Clément, *Swimming Bacteria in Poiseuille Flow: The Quest for Active Bretherton-Jeffery Trajectories*, *Europhys. Lett.* **126**, 44003 (2019).
- [30] M. Molaie, M. Barry, R. Stocker, and J. Sheng, *Failed Escape: Solid Surfaces Prevent Tumbling of Escherichia coli*, *Phys. Rev. Lett.* **113**, 068103 (2014).
- [31] L. Turner, L. Ping, M. Neubauer, and H. C. Berg, *Visualizing Flagella while Tracking Bacteria*, *Biophys. J.* **111**, 630 (2016).
- [32] P. S. Lovely and F. W. Dahlquist, *Statistical Measures of Bacterial Motility and Chemotaxis*, *J. Theor. Biol.* **50**, 477 (1975).
- [33] K. M. Taute, S. Gude, S. J. Tans, and T. S. Shimizu, *High-Throughput 3D Tracking of Bacteria on a Standard Phase Contrast Microscope*, *Nat. Commun.* **6**, 8776 (2015).
- [34] F. Perrin, *Mouvement Brownien d’un Ellipsoïde (I). Dispersion Diélectrique pour des Molécules Ellipsoïdales*, *J. Phys. Radium* **5**, 497 (1934).
- [35] F. Perrin, *Mouvement Brownien d’un Ellipsoïde (II). Rotation Libre et Dépolarisation des Fluorescences. Translation et Diffusion de Molécules Ellipsoïdales*, *J. Phys. Radium* **7**, 1 (1936).
- [36] N. Figueroa-Morales, *Active Bacterial Suspensions: From Microhydrodynamics to Transport Properties in Microfluidic Channels*, Ph. D. thesis, UPMC, Paris, 2016.
- [37] W. K. Smits, O. P. Kuipers, and J.-W. Veening, *Phenotypic Variation in Bacteria: The Role of Feedback Regulation*, *Nat. Rev. Microbiol.* **4**, 259 (2006).
- [38] A. J. Waite, N. W. Frankel, and T. Emonet, *Behavioral Variability and Phenotypic Diversity in Bacterial Chemotaxis*, *Annu. Rev. Biophys.* **47**, 595 (2018), pMID 29618219.
- [39] M. Wu, J. W. Roberts, S. Kim, D. L. Koch, and M. P. DeLisi, *Collective Bacterial Dynamics Revealed Using a Three-Dimensional Population-Scale Defocused Particle Tracking Technique*, *Appl. Environ. Microbiol.* **72**, 4987 (2006).
- [40] S. Y. S. Dufour, S. Gillet, S. Frankel, D. B. Weibel, and T. Emonet, *Direct Correlation between Motile Behavior and Protein Abundance in Single Cells*, *PLoS Comput. Biol.* **12**, e1005041 (2016).
- [41] S. Khan and R. M. Macnab, *The Steady-State Counter-clockwise/Clockwise Ratio of Bacterial Flagellar Motors Is Regulated by Protonmotive Force*, *J. Mol. Biol.* **138**, 563 (1980).
- [42] P. Cluzel, M. Surette, and S. Leibler, *An Ultrasensitive Bacterial Motor Revealed by Monitoring Signaling Proteins in Single Cells*, *Science* **287**, 1652 (2000).
- [43] E. Altshuler, G. Miño, C. Pérez-Penichet, L. del Río, A. Lindner, A. Rousselet, and E. Clément, *Flow-Controlled Densification and Anomalous Dispersion of E. coli through a Constriction*, *Soft Matter* **9**, 1864 (2013).
- [44] N. Figueroa-Morales, E. Altshuler, A. Hernández-García, A. Lage-Castellanos, and E. Clément, *Two-Dimensional Continuous Model for Bacterial Flows through Microfluidic Channels*, *Rev. Cubana Fis.* **30**, 3 (2013).
- [45] N. Figueroa-Morales, G. L. Miño, A. Rivera, R. Caballero, E. Clément, E. Altshuler, and A. Lindner, *Living on the Edge: Transfer and Traffic of E. coli in a Confined Flow*, *Soft Matter* **11**, 6284 (2015).
- [46] A. J. T. M. Mathijssen, N. Figueroa-Morales, G. Junot, É. Clément, A. Lindner, and A. Zöttl, *Oscillatory Surface Rheotaxis of Swimming E. coli Bacteria*, *Nat. Commun.* **10**, 3434 (2019).
- [47] G. Ariel, A. Rabani, S. Benisty, J. D. Partridge, R. M. Harshey, and A. Be’er, *Swarming Bacteria Migrate by Lévy Walk*, *Nat. Commun.* **6**, 8396 (2015).
- [48] S. Bianchi, F. Saglimbeni, and R. Di Leonardo, *Holographic Imaging Reveals the Mechanism of Wall Entrapment in Swimming Bacteria*, *Phys. Rev. X* **7**, 011010 (2017).
- [49] N. C. Darnton, L. Turner, S. Rojevsky, and H. C. Berg, *On Torque and Tumbling in Swimming Escherichia coli*, *J. Bacteriol.* **189**, 1756 (2007).
- [50] P. J. Mears, S. Koirala, C. V. Rao, I. Golding, and Y. R. Chemla, *Escherichia coli Swimming Is Robust against Variations in Flagellar Number*, *eLife* **3**, e01916 (2014).
- [51] J. Saragosti, P. Silberzan, and A. Buguin, *Modeling E. coli Tumbles by Rotational Diffusion. Implications for Chemotaxis*, *PLoS One* **7**, e35412 (2012).
- [52] D. Saintillan, *The Dilute Rheology of Swimming Suspensions: A Simple Kinetic Model*, *Exp. Mech.* **50**, 1275 (2010).
- [53] R. Soto, *Kinetic Theory and Transport Phenomena* (Oxford University, New York, 2016), Vol. 25.